

Short-Acting Narcotic Analgesics Review Therapeutic Class Review (TCR)

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FDA-APPROVED INDICATIONS

Drug	Federal	Manufacturer	Indication(s)
	Schedule		(c)
butorphanol nasal spray ¹	CIV	generic	Management of pain when the use of an opioid analgesic is appropriate
codeine sulfate ²	CII	generic	Mild to moderately severe pain
codeine/acetaminophen ³ (Tylenol® #3, Tylenol #4, Capital®)	CIII	generic	Mild to moderate pain
codeine/butalbital/aspirin/caffeine (Fiorinal® with codeine) ⁴	CIII	generic, Actavis	Tension or muscle contraction headache
dihydrocodeine bitartrate/ acetaminophen/caffeine (Trezix®) ⁵	CIII	generic, WraSer	Moderate to moderately severe pain
dihydrocodeine bitartrate aspirin/caffeine (Synalgos DC®)6	CIII	Caraco	Moderate to moderately severe pain
fentanyl buccal (Fentora®) ⁷	CII	Cephalon	Breakthrough cancer pain in patients with malignancies who are already receiving and who
fentanyl nasal spray (Lazanda®) ⁸	CII	Depomed	are tolerant to opioid therapy for their underlying persistent cancer pain
fentanyl sublingual spray (Subsys®) ⁹	CII	Insys	
fentanyl sublingual tablet (Abstral®) ¹⁰	CII	Galena Biopharma	
fentanyl transmucosal oral lozenge (Actiq®) ¹¹	CII	generic, Cephalon	
hydrocodone/acetaminophen solution (Hycet®, Lortab®, Zamicet®) ^{12,13,14}	CII	generic	Moderate to moderately severe pain
hydrocodone/acetaminophen tablet (Lorcet®, Lortab, Norco®, Verdrocet™, Vicodin®, Xodol®) ^{15,16}	CII	generic	
hydrocodone/ibuprofen (Ibudone®, Reprexain™, Vicoprofen®, Xylon™) ^{17,18}	CII	generic	Short-term management of acute pain
hydromorphone (Dilaudid®) ¹⁹	CII	generic, Purdue	Management of pain in patients where an opioid analgesic is appropriate
levorphanol ²⁰	CII	Sentynl	Moderate to severe pain
meperidine (Demerol®) ^{21*}	CII	generic	Moderate to severe pain
morphine immediate-release ²²	CII	generic	Moderate to severe acute and chronic pain
oxycodone immediate-release (Oxaydo™) ^{23†}	CII	Egalet	Moderate to severe acute and chronic pain
oxycodone (Roxicodone™) ^{24,25}	CII	generic	Moderate to severe pain



FDA-Approved Indications (continued)

Drug	Federal Schedule	Manufacturer	Indication(s)
oxycodone/acetaminophen (Endocet®, Percocet®, Primlev™, Roxicet™) ^{26,27}	CII	generic	Moderate to severe pain
oxycodone/acetaminophen (Xartemis™ XR)28	CII	Mallinckrodt	Management of acute pain severe enough to require opioid treatment
oxycodone/aspirin (Endodan®, Percodan®) ²⁹	CII	generic	Moderate to severe pain
oxycodone/ibuprofen ³⁰	CII	generic	Short-term (7 days or less) treatment of acute, moderate to severe pain
oxymorphone immediate-release (Opana®) ³¹	CII	generic	Moderate to severe acute pain
pentazocine/naloxone ³²	CIV	generic	Moderate to severe pain
tapentadol (Nucynta®) ³³	CII	Janssen/Depomed	Relief of moderate to severe acute pain
tramadol (Ultram®) ³⁴		generic	Management of moderate to moderately severe pain in adults
tramadol/acetaminophen (Ultracet®) ³⁵		generic	Short-term (5 days or less) treatment of acute pain

^{*} Meperidine should only be used for the acute treatment of moderate to severe pain. It should not be used for the treatment of chronic pain. Prolonged use can increase the risk of toxicity (e.g., seizures) from the accumulation of the metabolite, normeperidine.

OVERVIEW

Pain is often under-treated, and pain management is greatly misunderstood. Different management techniques are utilized for acute and chronic pain. It has been cited in studies that up to 73% of hospitalized medical patients receiving opiates were found in severe or moderate distress despite their analgesic regimen.³⁶ Caregivers' misconceptions regarding opiate doses, duration of analgesic effect, and fear of addiction were partly responsible for this under-treatment in both hospital and ambulatory care settings.³⁷

The World Health Organization's (WHO) guidelines for cancer pain management recommend a 3-stepped approach with consideration for the type of pain and response to therapy. Therapy for mild pain should include non-opioid analgesics, such as acetaminophen or non-steroidal anti-inflammatory drugs (NSAIDs). For mild to moderate pain, oral combinations of acetaminophen and NSAIDs with opioids are recommended. For moderate to severe pain, opioid analgesics are the mainstay.

The American Pain Society does not distinguish amongst the available products in their 2009 clinical guidelines for the use of chronic opioid therapy for the treatment of chronic non-cancer pain.⁴⁰ Titration of dose and frequency should be individualized to the patient's response and experience of



[†] Product name changed from Oxecta™ to Oxaydo.

adverse effects. In 2016, the American Pain Society published a guideline on the management of postoperative pain. ⁴¹ These guidelines recommend oral over intravenous opioid analgesics in patients who are able to use the oral route. Intramuscular opioids are not recommended. They also recommend multimodal pain control, including non-pharmacologic and other medications, such as acetaminophen or NSAIDs, gabapentin, or pregabalin; however, one opioid agent is not recommended over another.

In 2016, the Centers for Disease Control and Prevention (CDC) released guidelines for prescribing opioids for chronic pain outside of active cancer, palliative, and end-of-life care. 42 These guidelines include 12 recommendations: 3 regarding when to initiate or continue opioids for chronic pain; 4 regarding opioid selection, dosage, duration, follow-up, and discontinuation; and 5 regarding assessing risk and addressing harms of opioid abuse. The guidelines prefer nonpharmacologic and nonopioid pharmacologic therapy for chronic pain, and recommend a full individual assessment, including risk evaluation and realistic treatment goal setting, prior to prescribing opioids for chronic pain. If opioids are deemed appropriate for a patient's chronic pain, they recommend initial treatment with immediate-release opioids instead of extended-release opioids, which should be prescribed at the lowest effective dose. They further specify that doses of \geq 50 morphine milligram equivalents (MME)/day should prompt reassessment of the individual's benefits and risks and use of ≥ 90 MME/day should be avoided without justification. They state that long-term opioid use often begins with acute pain treatment; thus, opioids for acute pain should be immediate-release, the lowest effective dose, and the quantity should not exceed the expected duration of pain severe enough to require opioids (typically 3 days and with > 7 days rarely needed). They recommend reassessment within 1 to 4 weeks to determine benefits, harms, and appropriate dosing and continued follow up at least every 3 months. At these visits, efforts should be made to optimize other therapies and taper or discontinue opioids as able and as risks outweigh the individual's benefits. In order to decrease risks, the guidelines recommend avoiding concurrent use of benzodiazepines when possible and risk management strategies, such as offering naloxone in high-risk individuals (e.g., history of overdose, history of substance abuse, doses ≥ 50 MME/day, concurrent benzodiazepine use). Likewise, they recommend urine drug testing at baseline and annually and review of state prescription drug monitoring programs (PDMPs) at baseline and every 3 months. Prescribers should also offer treatment for opioid use disorder (e.g., buprenorphine or methadone in combination with behavioral therapies).

Over the years, various products within this class have been removed from the market, reformulated, or rescheduled based on abuse potential. In 2009, A Federal Drug Administration (FDA) Advisory Committee in 2009 recommended that all propoxyphene-containing products be removed from the market based on their low benefit-to-risk ratio, and this was enforced in 2010.⁴³

The FDA announced prescription acetaminophen combinations would be limited to a maximum of 325 mg acetaminophen per dosage unit. The FDA issued reminders for providers to stop prescribing/dispensing prescription combination products that contain acetaminophen 325 mg per tablet, capsule, or other dosage unit. These products are no longer considered safe by FDA and have been voluntarily withdrawn.⁴⁴

Hydrocodone combination products were under review for possible Federal Schedule promotion from III to II. In early December 2013, the FDA submitted a formal recommendation to the Department of Health and Human Services to make the change. The Drug Enforcement Agency (DEA) made its final decision regarding appropriate scheduling of hydrocodone-containing productions, resulting in the



reclassification of hydrocodone combination products from CIII to CII which took effect on October 6, 2014.⁴⁵

Soon after its approval in the United States (U.S.) in 1995, diversion and abuse of tramadol were reported. This led to the addition of warnings regarding the abuse potential of tramadol to the product labeling by the Food and Drug Administration (FDA). Tolerance, dependence and addiction to tramadol have been demonstrated and abrupt discontinuation of the drug can result in withdrawal symptoms. Effective August 18, 2014, tramadol-containing products were placed into Schedule IV of the Controlled Substance Act. 46

In April 2015, the FDA issued final guidance on the evaluation and labeling of abuse-deterrent opioids for industry. The only agent within this therapeutic class with abuse-deterrent properties is Oxaydo, and its labeling includes data from an abuse-deterrence study. The clinical significance of decreased "drug-liking" evaluated in the study, however, is not established.

In response to opioid abuse, the FDA announced an action plan in 2016 to approach opioid medications. The action plan includes an evaluation of risks and benefits, using experts to determine abuse-deterrence, and improving access to abuse-deterrent formulations and medication-assisted treatment options. In March 2016, the FDA announced that all immediate-release opioid pain medications would now require a new box warning about the serious risks of misuse, abuse, addiction, overdose, and death. 51

PHARMACOLOGY^{52,53,54,55,56,57,58,59,60,61,62,63,64,65,66,67,68,69,70,71,72,73,74,75,76,77,78}

Opioid agonists reduce pain by acting primarily through interaction with opioid mu-receptors located in the brain, spinal cord, and smooth muscle. The primary site of therapeutic action is the central nervous system (CNS). Stimulation at this receptor produces supraspinal analgesia, respiratory depression, euphoria, and physical dependence. Opioid agonists produce respiratory depression by direct action on the brain stem respiratory center.

The opioid agents in this review can be divided into full agonists and mixed agonist/antagonists. The weaker full agonists, such as hydrocodone, codeine, and tramadol, are often prescribed in combination with nonopioid analgesics. Strong full agonists, such as fentanyl, meperidine, morphine, hydromorphone, oxymorphone, levorphanol, and oxycodone, are generally used for treatment of moderate to severe pain.

Butorphanol and pentazocine are mixed agonist-antagonist agents. They are both weak antagonists at μ -receptors and agonists at kappa-receptors. Due to their action at the kappa-receptors, these agents may produce dysphoric effects and increased blood pressure and heart rate in some individuals. Due to their opioid antagonist properties, there is a ceiling on the analgesic effects of pentazocine and butorphanol.

Tramadol (Rybix ODT, Ultram, and Ultracet) are centrally-acting analgesics with dual opioid and nonopioid mechanisms. In addition to activity at opioid receptors, tapentadol (Nucynta) inhibits norepinephrine re-uptake and tramadol weakly inhibits norepinephrine and serotonin re-uptake.

Aspirin and NSAIDs work by blocking cyclooxygenase (COX)-1 and COX-2, which prevent the synthesis of various prostaglandins. These prostaglandins are partially responsible for the development of pain and inflammation.



The exact mechanism of action for acetaminophen is unknown, but it mediates its actions centrally. Acetaminophen is thought to act primarily in the CNS and increases the pain threshold by inhibiting COX-1 and COX-2. Unlike NSAIDs, acetaminophen does not inhibit COX in peripheral tissues. Acetaminophen may also decrease sensitization of pain receptors to mechanical or chemical stimulation.

Caffeine causes cerebral vasoconstriction, which decreases blood flow and oxygen tension. In combination with acetaminophen, caffeine may provide a quicker onset of action and enhance pain relief allowing for lower doses of analgesics.

Naloxone, an opioid antagonist, has no pharmacologic activity when administered orally at 0.5 mg. Studies in animals indicate that the presence of naloxone does not affect pentazocine analgesia when the combination is given orally. If the combination is given by injection, the action of pentazocine is neutralized.



PHARMACOKINETICS^{79,80,81,82,83,84,85,86,87,88,89,90,91,92,93,94,95,96,97,98,99,100,101,102,103,104,} 105

Drug	Half-Life (hr)	Tmax (hr)	Excretion
- 0		id Component	
butorphanol nasal spray	4.7 – 6.6 (parent) 18 (metabolite)	0.6 – 1	extensively metabolized and excreted in urine and feces
codeine sulfate	3 – 4 (parent) 2 (metabolite – morphine)	No data available	primarily urine
dihydrocodeine ¹⁰⁶	3.3 – 4.5	No data available	metabolized to active dihydromorphine and renally eliminated
fentanyl buccal (Fentora)	2.63 – 11.7	0.5 – 0.75	>90% metabolized and renally eliminated
fentanyl nasal spray (Lazanda)	15.0 – 24.9	0.25 - 0.35	primarily (>90%) eliminated by biotransformation to N-dealkylated and hydroxylated inactive metabolites.
fentanyl sublingual spray (Subsys)	5.25 – 11.99	0.67 – 1.25	primarily (>90%) eliminated by biotransformation to N-dealkylated and hydroxylated inactive metabolites
fentanyl sublingual tablet(Abstral)	5 – 13.5	0.5 – 1	>90% metabolized and renally eliminated
fentanyl transmucosal oral lozenge (Actiq)	3.2 – 6.4	0.33 – 0.67	>90% metabolized and renally eliminated
hydrocodone	3.8	1.3 – 3	hydrocodone and metabolites renally eliminated
hydromorphone (Dilaudid)	2.3	0.73	highly metabolized
levorphanol	11 – 16	1	extensively metabolized and renally eliminated
meperidine (Demerol)	3 – 4 (parent) 15 – 30 (metabolite)	2	highly metabolized and renally eliminated
morphine immediate- release	2 – 15	0.5	extensively metabolized and renally eliminated
oxycodone	3 – 4	1.2 – 1.4	primarily metabolized and renally eliminated
oxycodone/acetaminophe n (Xartemis XR)	oxycodone 4.5	oxycodone 3 – 4	oxycodone: primarily metabolized and renally eliminated; acetaminophen: primary hepatic metabolism and primarily eliminated by glucuronide and sulfate conjugates
oxymorphone immediate- release (Opana)	7.3 – 9.4	No data available	highly metabolized and eliminated in urine and feces
pentazocine	0.5 – 4	3.6	extensively metabolized and renally eliminated
tapentadol (Nucynta)	4	1.25	highly metabolized eliminated in urine



Pharmacokinetics (continued)

Drug	Half-Life (hr)	Tmax (hr)	Excretion				
	Opioid Component						
tramadol	6.3 (tramadol) 7.4 (metabolites)	2-3	60% metabolized to active metabolites				
Drug	Half-Life (hr)	Tmax (hr)	Excretion				
	Non-opioid Component						
butalbital	35	No data available	metabolized eliminated in urine				
acetaminophen acetaminophen (Xartemis XR)	1 – 3 5.8	1.2 – 3 0.75–1	highly metabolized and renally eliminated				
aspirin	0.25 – 0.3	2 – 3 (low dose) 15 – 30 (high dose)	highly metabolized and renally eliminated				
caffeine ¹⁰⁷	No data available	3	highly metabolized and renally eliminated				
ibuprofen	1.8 – 2.6	1.6 – 3.1	highly metabolized renally excreted				
naloxone	2-3	1-3	highly metabolized and renally eliminated				

CONTRAINDICATIONS/WARNINGS^{108,109,110,111,112,113,114,115,116,117,118,119,120,121,} 122,123,124,125,126,127,128,129,130,131,132,133,134

All immediate-release opioid pain medications contain a box warning regarding serious risks of misuse, abuse, addiction, overdose, and death. 135

These agents are contraindicated in patients with known hypersensitivity to opioids or other components of the product. Patients known to be hypersensitive to opioids may exhibit cross sensitivity in the class. Hydromorphone liquid formulation contains sodium metabisulfite which may cause allergic-type reactions in susceptible patients.

In general, opioids are contraindicated in patients who have acute or severe bronchial asthma or hypercarbia; situations of significant respiratory depression (in the absence of resuscitative equipment or monitors); and patients with known or suspected paralytic ileus. Agents containing butorphanol, hydrocodone, levorphanol, meperidine, and pentazocine do not list these conditions as contraindications, but warnings to use with caution if any of the conditions are present.

Hydromorphone (Dilaudid) liquid and 8 mg tablets are contraindicated in patients for obstetrical analgesia.

Opioids should be used with caution in patients with renal or hepatic impairment and dosage adjustments may be warranted depending on the specific agent and degree of impairment. Oxymorphone is contraindicated in patients with moderate or severe hepatic impairment.

Monoamine oxidase inhibitors (MAOI) can markedly potentiate the action of opioid agents; therefore, opioid use is not recommended in patients currently taking MAOIs or within the previous 14 days. Caution should be observed in administering pentazocine to patients who are currently receiving MAOIs or who have received them within the preceding 14 days, due to potential CNS excitation and hypertension due to catecholamines effects. In addition, all opioids contain a warning regarding



serotonin syndrome when used concomitantly with any serotonergic drug (e.g., MAO inhibitors, selective serotonin reuptake inhibitors (SSRIs), serotonin and norepinephrine reuptake inhibitors (SNRI), tricyclic antidepressants (TCAs), triptans, linezolid, or lithium). Serotonin syndrome typically occurs within several hours to a few days following use.¹³⁶

Opioids may induce or aggravate seizures in some clinical settings, particularly in those patients with a history of seizure disorders.

Tramadol-containing products are contraindicated in any situation where opioids are contraindicated including acute intoxication with any of the following: alcohol, hypnotics, narcotics, centrally-acting analgesics, opioids, or psychotropic drugs. Tramadol may worsen central nervous system and respiratory depression in these patients.

Oxycodone/ibuprofen and hydrocodone/ibuprofen (Ibudone, Reprexain, Vicoprofen/ES/HP, and Xylon) are contraindicated in the treatment of peri-operative pain in the coronary artery bypass graft (CABG) setting. The black box warnings for NSAID-containing products cite the increased risk for adverse events seen with NSAID use, such as serious cardiovascular thrombotic events, myocardial infarction, stroke, and gastrointestinal adverse events, all of which can be fatal.

Tramadol (Ultram) and tramadol/acetaminophen (Ultracet) are contraindicated in any situation where opioids are contraindicated including acute intoxication with any of the following: alcohol, hypnotics, narcotics, centrally-acting analgesics, opioids, or psychotropic drugs. Tramadol may worsen CNS and respiratory depression in these patients. Withdrawal symptoms may occur if tramadol is discontinued abruptly. Clinical experience suggests that withdrawal symptoms may be avoided by tapering tramadol at the time of discontinuation.

Acetaminophen/caffeine/dihydrocodeine (Trezix) is contraindicated in patients with hypersensitivity to any of the components or in situations where opioids are contraindicated. These include significant respiratory depression, particularly in unmonitored settings or in the absence of resuscitation equipment, acute or severe bronchial asthma, hypercapnia, or paralytic ileus.

Respiratory depression and death have occurred in children with obstructive sleep apnea who received codeine in the post-operative period following tonsillectomy and/or adenoidectomy and had evidence of being ultra-rapid metabolizers of codeine. Codeine-containing products, including aspirin/caffeine/dihydrocodeine (Synalgos DC), are contraindicated for post-operative pain management in all pediatric patients undergoing tonsillectomy and/or adenoidectomy.

All products in this class should be used with caution in patients who may be susceptible to intracranial effects of carbon dioxide retention, such as those with evidence of increased intracranial pressure or impaired consciousness. Opioids may obscure the clinical course of a patient with a head injury and should be employed only if clinically warranted.

Opioids produce peripheral vasodilation which may result in orthostatic hypotension for some patients. Additionally, gastrointestinal opioid-induced effects may include a reduction in gastric, biliary, and pancreatic secretions.

Butorphanol and pentazocine can elevate blood pressure and heart rate. Particular caution should be exercised in conditions where alterations in vascular resistance and blood pressure might be particularly undesirable, such as in the acute phase of myocardial infarction.



Meperidine should be used with caution in patients with atrial flutter or other supraventricular tachycardias due to a possible vagolytic action that may produce a significant increase in ventricular response rate.

Opioids depress the cough reflex by direct effect on the cough center in the medulla. Caution should be exercised in postoperative use and in patients with pulmonary disease.

Other warnings instruct prescribers to be aware of the abuse potential of these products, the possibility of hypoventilation, the dangers to pediatric patients if used, and the increased risk of respiratory depression when used with CYP450 3A4 inhibitors. Impairment of physical and/or mental abilities, increased seizure risk, use of caution when performing hazardous tasks, respiratory depression, abuse potential, and increased sedation when used with other central nervous system depressants are also associated with opioid use. Opioids diminish propulsive peristaltic waves in the gastrointestinal tract and may obscure the diagnosis or clinical course in patients with acute abdominal conditions. Monitor for decreased bowel motility.

Opiate agonists can cause urinary retention and oliguria due to increased tension of the detrusor muscle. Patients more prone to these effects include those with prostatic hypertrophy, urethral stricture, bladder obstruction, or pelvic tumors. Drug accumulation or prolonged duration of action can occur in patients with renal impairment. Fentanyl buccal (Fentora) contains a black box warning regarding abuse potential; while both fentanyl buccal and fentanyl sublingual (Subsys) include a black box warning citing risks of respiratory depression and, when dispensing, there should be no substitution of any other fentanyl products.

Opioids inhibit the secretion of adrenocorticotropic hormone (ACTH) and cortisol. All opioids carry a warning for adrenal insufficiency; if an opioid causes adrenal insufficiency, treat with corticosteroids and withdraw the opiate as appropriate. Also, thyroid stimulating hormone may be stimulated or inhibited by opioids. Patients with adrenal insufficiency, thyroid disease (e.g., hypothyroidism), or myxedema may not be appropriate candidates for codeine administration.

Patients receiving therapeutic doses of pentazocine/acetaminophen have experienced hallucinations (usually visual), disorientation, and confusion which have cleared spontaneously within a period of hours. Visual blurring, dysphoria, and hallucinations have been reported rarely with butorphanol. Hallucinations, suicidal ideation, and panic attack have been reported in after-market surveillance of tapentadol (Nucynta).

Particular caution should be exercised in administering pentazocine to patients with porphyria since it may provoke an acute attack in susceptible individuals.

Opioid analgesics may cause tolerance and/or physical dependence with chronic use. Withdrawal symptoms may occur if these agents are discontinued abruptly and may be avoided by tapering opioid dosage at the time of discontinuation.

Due to their opioid antagonist properties, pentazocine and butorphanol can precipitate withdrawal symptoms in patients physically dependent on full agonists. Such patients should have an adequate period of withdrawal from opioid drugs prior to beginning butorphanol therapy.

Fentanyl nasal spray (Lazanda) and fentanyl sublingual spray (Subsys) should not be used for acute or post-operative pain. On a microgram per microgram basis, fentanyl nasal spray and sublingual spray are not equivalent to any other fentanyl products due to differences in pharmacokinetics.



In 2009, an FDA Advisory Committee recommended that the FDA put more restrictions on acetaminophen use in an effort to curb overdoses that can cause liver failure and/or death. 138

In 2011, the FDA asked manufacturers of prescription acetaminophen combination products to limit the maximum amount of acetaminophen in these products to 325 mg per tablet, capsule, or other dosage unit by January 1, 2014. ¹³⁹

Aspirin is contraindicated in patients with known allergy to nonsteroidal anti-inflammatory drug products and in patients with the syndromes of asthma, rhinitis, and nasal polyps. Aspirin may cause severe urticaria, angioedema, or bronchospasm (asthma). Aspirin should not be used in children or teenagers for viral infections, with or without fever, because of the risk of Reye syndrome. Patients who consume 3 or more alcoholic drinks every day should be counseled about the bleeding risks involved with chronic, heavy alcohol use while taking aspirin. Even low doses of aspirin can inhibit platelet function leading to an increase in bleeding time. Patients with a history of active peptic ulcer disease should avoid using aspirin, which can cause gastric mucosal irritation and bleeding.

Risk Evaluation and Mitigation Strategy (REMS)¹⁴⁰

Due to the risk of misuse, abuse, addiction, and overdose related to transmucosal fentanyl formulations (fentanyl sublingual [Abstral], fentanyl oral transmucosal [Actiq], fentanyl buccal [Onsolis, Fentora], fentanyl nasal spray [Lazanda], and fentanyl sublingual spray [Subsys]), these agents are only available through a restricted access program called Transmucosal Immediate-release Fentanyl (TIRF) Risk Evaluation and Mitigation Strategy (REMS) Access Program. These medications are also dispensed with medication guides. Outpatient health providers including prescribers and pharmacies must enroll in this program. Wholesalers and distributors also must enroll in order and distribute only to authorized pharmacies. In addition, outpatients must sign a Patient-Prescriber Agreement to ensure they understand the risks and benefits of therapy.

DRUG INTERACTIONS^{141,142,143,144,145,146,147,148,149,150,151,152,153,154,155,156,157,158,} 159,160,161,162,163,164,165,166,167

All opioid agents should be used with caution and in reduced dosage in patients who are concurrently receiving other narcotic analgesics, muscle relaxants, general anesthetics, phenothiazines, other tranquilizers, sedative-hypnotics, tricyclic antidepressants, and other CNS depressants (including alcohol). Respiratory depression, hypotension, and profound sedation or coma may result.

Monoamine oxidase inhibitors (MAOIs) may intensify the actions of opioid agents. All opioids contain a warning regarding serotonin syndrome when used concomitantly with any serotonergic drug as described above. 168

Fentanyl and tramadol are mainly metabolized by the CYP450 enzyme pathway; co-administration of these agents with CYP450 enzyme inducers or inhibitors may adversely affect their metabolism. Patients taking carbamazepine may have a significantly reduced analgesic effect of tramadol; concurrent administration of carbamazepine and tramadol is not recommended due to the increased tramadol metabolism by carbamazepine and because of the seizure risk associated with tramadol.

Patients taking cytochrome CYPP450 enzyme inducers or inhibitors may demonstrate an altered response to codeine; therefore, analgesic activity should be monitored. Acyclovir may increase the plasma concentration of meperidine and normeperidine. Ritonavir may increase the plasma



concentration of normeperidine. Phenytoin may increase the metabolism and clearance of meperidine. Caution should be used with concomitant use of meperidine with any of these agents.

Concurrent use of medications with anticholinergic activity and opioid analgesics may result in increased risk of urinary retention and/or severe constipation and paralytic ileus.

CNS side effects (e.g., confusion, disorientation, respiratory depression, apnea, seizures) have been reported following co-administration of cimetidine with opioid analgesics; a causal relationship has not been established.

Agonist/antagonist analgesics (pentazocine, butorphanol) should be administered with caution to patients receiving a pure opioid agonist analgesic. In this situation, mixed agonist/antagonist analgesics may reduce the analgesic effect of the full opioid agonist and/or may precipitate withdrawal symptoms in these patients.

A slower onset can be anticipated if butorphanol tartrate nasal spray is administered concomitantly with, or immediately following, a nasal vasoconstrictor due to a decreased rate of absorption.

Co-administration of a vasoconstrictive nasal decongestant, such as oxymetazoline, to treat allergic rhinitis leads to lower peak plasma concentrations and a delayed Tmax of fentanyl that may cause fentanyl nasal spray (Lazanda) to be less effective in patients with allergic rhinitis who use such decongestants, thus potentially impairing pain management.

Due to the ibuprofen component, hydrocodone/ibuprofen and oxycodone/ibuprofen are associated with interactions with ACE inhibitors, methotrexate, and warfarin that are more frequently seen with NSAID co-administration. Ibuprofen has been shown to reduce the natriuretic effect of furosemide and thiazides in some patients. Ibuprofen has been shown to elevate plasma lithium concentration and reduce renal lithium clearance.

Chronic and excessive consumption of alcohol may increase the hepatotoxic risk of acetaminophen. The potential for hepatotoxicity with acetaminophen also may be increased in patients receiving anticonvulsants that induce hepatic microsomal enzymes (including phenytoin, barbiturates, and carbamazepine) or isoniazid.

Aspirin may enhance the effects of anticoagulants and inhibit the uricosuric effects of uricosuric agents.

Caffeine may enhance the cardiac inotropic effects of beta-adrenergic stimulating agents. Co-administration of caffeine and disulfiram may lead to a substantial decrease in caffeine clearance. Caffeine may increase the metabolism of other drugs, such as phenobarbital and aspirin. Caffeine accumulation may occur when products or foods containing caffeine are consumed concomitantly with quinolones, such as ciprofloxacin.



ADVERSE EFFECTS 169,170,171,172,173,174,175,176,177,178,179,180,181,182,183,184,185,186,187,188,189,190,191,192,193,194,195

Drug	Asthenia	Constipation	Dizziness	Dyspnea	Headache	Nausea	Rash	Somnolence	Vomiting
butorphanol NS	>1	>1	19	> 1	>1	≤ 13	>1	43	≤ 13
codeine sulfate	nr	reported	reported	nr	reported	reported	reported	reported	reported
codeine/acetaminophen (Tylenol #3, Tylenol #4, Capital)	nr	reported	reported	reported	nr	reported	reported	nr	reported
codeine/butalbital/aspirin/ caffeine (Fiorinal with codeine)	nr	nr	reported	nr	nr	reported	reported	reported	reported
dihydrocodeine bitartrate/ acetaminophen/caffeine (Trezix)	nr	reported	reported	nr	nr	reported	nr	reported	reported
dihydrocodeine bitartrate/ aspirin/caffeine (Synalgos DC)	nr	reported	reported	nr	nr	reported	nr	reported	reported
fentanyl buccal (Fentora)	11	12	13 – 19	9	9 – 10	17 – 29	< 1	7 – 9	5 – 20
fentanyl nasal spray (Lazanda)	≥1	1-10	≥1	≥ 1	≥1	4 – 9	nr	≥ 1	7 – 13
fentanyl sublingual spray (Subsys)	9.7	5 – 10.4	7.2	10.4	≥1	10.4 – 13.1	nr	9.5	10.3 – 16
fentanyl sublingual tablet (Abstral)	reported	4.8	reported	0.6	3	6	reported	reported	reported
fentanyl transmucosal oral lozenge (Actiq)	9 – 38	4 – 20	16 – 17	4 – 22	6 – 20	23 – 45	2-8	15 – 17	12 – 31
hydrocodone/ acetaminophen solution (Hycet, Lortab, Zamicet)	nr	reported	reported	reported	nr	reported	reported	reported	reported
hydrocodone/ acetaminophen tablet Lorcet, Lortab, Norco, Verdrocet, Vicodin, Xodol)	nr	reported	reported	reported	nr	reported	reported	reported	reported



Adverse Effects (continued)

Drug	Asthenia	Constipation	Dizziness	Dyspnea	Headache	Nausea	Rash	Somnolence	Vomiting
hydrocodone/ibuprofen (Ibudone, Reprexain, Vicoprofen, Xylon)	3 – 9	22	14	< 3	27	21	<1	22	3 – 9
hydromorphone (Dilaudid)	reported	reported	reported	reported	reported	reported	reported	reported	reported
levorphanol	nr	nr	reported	nr	nr	reported	reported	nr	reported
meperidine (Demerol)	reported	reported	reported	nr	reported	reported	reported	nr	reported
morphine immediate- release	nr	reported	reported	nr	reported	reported	reported	reported	reported
oxycodone immediate- release (Oxaydo)	reported	reported	reported	reported	reported	reported	reported	reported	reported
oxycodone immediate- release (Roxicodone)	≥3	≥ 3	≥ 3	< 3	≥3	≥ 3	< 3	≥ 3	≥ 3
oxycodone/acetaminophen (Endocet, Percocet, Primlev, Roxicet)	reported	reported	reported	reported	reported	reported	reported	reported	reported
oxycodone/acetaminophen (Xartemis XR)	reported	reported	reported	reported	reported	reported	reported	reported	reported
oxycodone/aspirin (Endodan, Percodan)	reported	reported	reported	reported	reported	reported	reported	reported	reported
oxycodone/ibuprofen	3.3	4.5	5.1 – 19.2	< 1	10.2	8.8 – 25.4	< 2	7.3 – 17.4	4.5 – 5.3
oxymorphone immediate- release (Opana)	< 1	4	7	< 1	7	19	< 1	9	9
pentazocine/naloxone	reported	reported	reported	nr	reported	reported	reported	reported	reported
tapentadol (Nucynta)	nr	8	24	< 1	reported	30	1	15	18
tramadol (Ultram)	6 – 12	24 – 46	26 – 33	< 1	18 – 32	24 – 40	1-<5	16 – 25	9 – 17
tramadol/ acetaminophen (Ultracet)	>1	6	3	< 1	>1	3	>1	6	>1

Adverse effects are reported as a percentage. Adverse effects data are obtained from package inserts and are not meant to be comparative or all inclusive. nr = not reported.



Opioids have been associated with a decrease in sex hormone levels. Laboratory assessment is recommended in patients who report low libido, impotence, erectile dysfunction, lack of menstruation, or infertility. 196

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Pediatrics

Fentanyl buccal (Fentora), sublingual (Abstral), fentanyl nasal spray (Lazanda), fentanyl sublingual spray (Subsys), and tapentadol (Nucynta) are indicated for patients 18 years of age or older. Fentanyl transmucosal (Actiq) is approved for patients 16 years old or older. The safety and efficacy of tramadol-containing (Ultracet and Ultram) products in children under 16 years of age have not been studied, and their use is not recommended. Hydrocodone/ibuprofen (Ibudone, Reprexain, Vicoprofen, Xylon) has no established safety and efficacy in patients less than 16 years of age. Oxycodone/ibuprofen is safe and effective in patients 14 years and older. The safety and efficacy of pentazocine-containing products in children under 12 years of age have not been established. Hydrocodone/acetaminophen (Hycet and Lortab Elixir only) has not been studied in patients younger than two years old. Hydrocodone/acetaminophen (Lorcet, Norco, Vicodin/ES/HP) has not been adequately studied in pediatric patients. The safety and efficacy of the remaining products in this review have not been established in the pediatric population.

Pregnancy

The products listed in this review are Pregnancy Category C, except oxycodone single-ingredient products (Oxaydo, Roxicodone) which are Category B, and oxycodone/ibuprofen which is Pregnancy Category C prior to 30 weeks gestation; Category D starting at 30 weeks gestation.

Geriatrics

Opioid products should be used with caution in elderly patients due to greater sensitivity of primary effects and adverse effects. Doses should be titrated to provide adequate efficacy while minimizing risk.

Plasma levels of oxymorphone may be seen up to 40% higher in elderly patients over age 65 years than seen in younger patients. For elderly patients over 75 years old, total tramadol dose should not exceed 300 mg/day.

In the 2009 Management of Persistent Pain in Older Persons guideline, the American Geriatric Society (AGS) advises that in the elderly even pain that is causing severe impairment may not be spontaneously revealed for a variety of personal, cultural, or psychological reasons. Older persons may under-report pain, but there are also inherent difficulties in recognizing pain experienced by patients with cognitive impairment.²²⁴ However, all patients with moderate to severe pain, pain-related functional impairment, or diminished quality of life due to pain should be considered for opioid therapy and should be reassessed for ongoing attainment of therapeutic goals, adverse effects, and safe and responsible medication use. Tramadol has opioid activity with apparently low abuse potential and is reportedly about as effective and safe as codeine or hydrocodone. However, tramadol has the additional low risk of inducing seizures.



Hepatic and Renal Impairment

All agents in this review should be used with caution in patients with hepatic or renal impairment. Dosage reductions may be warranted.

Oxymorphone is contraindicated in patients with moderate to severe hepatic impairment.

Tapentadol should be used with caution in patients with moderate hepatic impairment. Patients with severe renal or hepatic impairment should not use tapentadol.

Tramadol should be given every 12 hours for patients with creatinine clearance (CrCl) < 30 mL/minute with a maximum dose of 200 mg per day. Patients with cirrhosis should receive tramadol 50 mg every 12 hours.

Other

Some individuals may be ultra-rapid metabolizers of codeine due to a specific cytochrome P450 2D6 (CYP2D6) phenotype and may convert codeine into its active metabolite, morphine, more rapidly and completely resulting in higher than expected serum morphine levels. Even at labeled dosage regimens, individuals who are ultra-rapid metabolizers may experience overdose symptoms such as extreme sleepiness, confusion, or shallow breathing.

The prevalence of this CYP2D6 phenotype varies widely and has been estimated at 0.5% to 1% in Chinese and Japanese, 0.5% to 1% in Hispanics, 1% to 10% in Caucasians, 3%% in African Americans, and 16% to 28% in North Africans, Ethiopians, and Arabs. Data are not available for other ethnic groups.

Cardiac Disease

Fentanyl buccal, sublingual tablet/spray, transmucosal, and nasal spray should be used with caution in patients with bradyarrhythmias.



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Drug	Starting Dose	Dosing Instructions	Available Strengths
butorphanol nasal spray	1 spray into 1 or both nostrils; may repeat after 3 to 4 hours	If 1 spray is administered and adequate pain relief is not achieved within 60 to 90 minutes, an additional 1 mg dose may be given The initial 2-dose sequence may be repeated in 3 to 4 hours, as required, after the second dose of the sequence	Solution: 10 mg/mL
codeine sulfate	15 to 60 mg every 4 to 6 hours, as needed	Do not exceed 360 mg in 24 hours	Tablets: 15, 30, 60 mg
codeine/ acetaminophen (Tylenol #3, Tylenol #4, Capital)	Tablet: 1 to 2 every 4 hours, as needed Elixir: 15 mL every 4 hours	Do not exceed codeine 60 mg per dose and 360 mg per day or acetaminophen 4 g per day	Tablets: 15/300 mg, 30/300 mg, and 60/300 mg Elixir: 12/120 mg per 5 mL Suspension (Capital): 12/120 mg per 5 mL
codeine/butalbital/ aspirin/caffeine (Fiorinal with codeine)	1 to 2 capsules every 4 hours	Do not exceed 6 capsules per day	Capsule: codeine 30 mg/butalbital 50 mg/aspirin 325 mg/caffeine 40 mg
dihydrocodeine bitartrate/ acetaminophen/ caffeine (Trezix)	2 capsules every 4 hours, as needed	Do not exceed 10 capsules per 24 hours	Capsule: acetaminophen 320.5 mg/ caffeine 30 mg/ dihydrocodeine 16 mg
dihydrocodeine bitartrate aspirin/caffeine (Synalgos DC)	2 capsules every 4 hours, as needed	Do not exceed 10 capsules per 24 hours	Capsule: aspirin 356.4 mg/caffeine 30 mg/dihydrocodeine 16mg
fentanyl buccal (Fentora)	100 mcg, as needed	Until the appropriate dose is reached, patients may find it necessary to use an additional unit during a single episode of breakthrough pain not relieved in 30 minutes; One tablet of the same dose may be taken; If pain is not relieved, patients must wait 4 hours before treating another episode of breakthrough pain If treatment of several consecutive breakthrough cancer pain episodes requires more than 1 unit per episode, an increase in dose to the next higher available strength should be considered If patient is currently on fentanyl transmucosal lozenges (Actiq), see prescribing information for additional dosing recommendations	Tablets: 100, 200, 400, 600, 800 mcg



Drug	Starting Dose	Dosing Instructions	Available Strengths
fentanyl nasal spray (Lazanda)	100 mcg, as needed	Individually titrate to an effective dose, from 100 mcg to 200 mcg to 400 mcg, and up to a maximum of 800 mcg, that provides adequate analgesia with tolerable side effects Dose is a single spray into 1 nostril, a single spray into each nostril (2 sprays), or 2 sprays into each nostril (4 sprays); no more than 4 doses per 24 hours Wait at least 2 hours before treating another episode of breakthrough pain with fentanyl nasal spray	Nasal sprays: 100, 400 mcg
fentanyl sublingual spray (Subsys)	100 mcg, as needed	Titrated as tolerated to an effective dose One dose of Subsys should be used per breakthrough pain episode; incases where the pain may not be relieved within 30 minutes of the dose, 1 additional dose of the same strength may be used for that breakthrough episode At least 4 hours must elapse prior to initiating treatment for another episode of pain Maintenance dosing should not exceed 4 doses per 24 hours Dose increase should be considered when several consecutive attempts to control breakthrough pain have failed	Sublingual sprays: 100, 200, 400, 600, 800, 1,200, 1,600 mcg
fentanyl sublingual tablet (Abstral)	100 mcg, as needed	Doses may be supplemented 1 time after 30 minutes; do not use more than 2 doses per episode of breakthrough pain; wait 2 hours before treating another episode Titrate to a successful dose and limit use to 4 episodes per day	Tablets: 100, 200, 300, 400, 600, 800 mcg
fentanyl transmucosal oral lozenge (Actiq)	200 mcg, as needed	Until the appropriate dose is reached, patients may find it necessary to use an additional unit during a single episode; patients must wait at least 4 hours before treating another episode of breakthrough pain If treatment of several consecutive breakthrough cancer pain episodes requires more than 1 unit per episode, an increase in dose to the next higher available strength should be considered	Transmucosal oral lozenges: 200, 400, 600, 800, 1,200, 1,600 mcg
hydrocodone/ acetaminophen solution (Hycet, Lortab, Zamicet)	15 mL every 4 to 6 hours	Not to exceed 90 mL in 24 hours See dosing chart in prescribing information for initial doses for children	Solution: 7.5/325 mg per 15 mL (Hycet), 10/300 mg per 15 mL (Lortab), 10/325 mg per 15 mL (Zamicet)



Drug	Starting Dose	Dosing Instructions	Available Strengths
hydrocodone/ acetaminophen tablet (Lorcet, Lortab, Norco, Verdrocet, Vicodin, Xodol)	1 to 2 tablets every 4 to 6 hours	Not to exceed 6 tablets or capsules in 24 hours; for tablets or capsules that contain 8 mg hydrocodone, may take up to 8 tablets per 24 hours For tablets that contain 7.5 or 10 mg hydrocodone, take 1 tablet or capsule every 4 to 6 hours	Tablets: 2.5/325 mg (Verdrocet), 5/300 mg (Vicodin, Xodol), 5/325 mg (Lorcet, Lortab, Norco), 7.5/300 mg (Vicodin ES, Xodol), 7.5/325 mg (Lorcet Plus, Lortab, Norco), 10/300 mg (Vicodin HP, Xodol), 10/325 mg (Lorcet HD, Lortab, Norco)
hydrocodone/ ibuprofen(Ibudone, Reprexain, Vicoprofen, Xylon)	1 tablet every 4 to 6 hours	Not to exceed a maximum of 5 tablets in 24 hours	Tablets: 2.5/200 mg (Reprexain), 5/200 mg (Ibudone, Reprexain), 7.5/200 mg (Vicoprofen), 10/200 mg (Ibudone, Reprexain, Xylon)
hydromorphone (Dilaudid)	Tablets: 2 to 8 mg every 4 to 6 hours Liquid: 2.5 to 10 mg every 3 to 6 hours	Dose should be adjusted so that at least 3 to 4 hours of pain relief may be achieved Dose should be increased, as needed, according to patient's response	Tablets: 2, 4, 8 mg Liquid: 5 mg/5 mL
levorphanol	2 mg every 6 to 8 hours	Total oral daily doses of more than 6 to 12 mg in 24 hours are generally not recommended as starting doses	Tablet: 2 mg
meperidine (Demerol)	Adult: 50 to 150 mg every 3 to 4 hours Pediatric: 1.1 to 1.8 mg/kg every 3 to 4 hours	Not for chronic use	Tablets: 50, 100 mg Solution: 50 mg/5 mL
morphine immediate-release	Tablets: 15 to 30 mg every 4 hours, as needed Solution: 10 to 20 mg every 4 hours, as needed	The dose should be titrated based upon the individual patient's response	Tablets: 15, 30 mg Solution: 10 mg/5 mL, 20 mg/5 mL, 100 mg/5 mL
oxycodone immediate-release (Oxaydo)	Opioid-naïve: 5 to 15 mg every 4 to 6 hours, as needed	The dose must be swallowed whole and is not amenable to crushing and dissolution Do not use for administration via nasogastric, gastric, or other feeding tubes as it may cause obstruction of the feeding tube	Tablet: 5, 7.5 mg (Abuse-deterrent; resistant to crushing, chewing, snorting, and injection related abuse)



Drug	Starting Dose	Dosing Instructions	Available Strengths
oxycodone immediate-release (Roxicodone)	5 to 15 mg every 4 to 6 hours, as needed	The dose should be titrated based upon the individual patient's response	Capsule: 5 mg Tablets: 5, 10, 15, 20, 30 mg Solution: 5 mg/5 mL, 20 mg/mL
oxycodone/ acetaminophen (Endocet, Percocet, Primlev, Roxicet)	1 to 2 tablets or capsules every 6 hours	Do not exceed oxycodone 60 mg or acetaminophen 4 g per day in adults Children: < 45 kg body weight – do not exceed 90 mg/kg per	Percocet and Endocet (tablets): 2.5/325, 5/325, 7.5/325, 10/325 mg
		day based on the acetaminophen component > 45 kg body weight – do not exceed 4 g per day	Primlev (tablets): 5/300, 7.5/300, 10/300 mg
		based on the acetaminophen component	Roxicet (tablet): 5/325 Roxicet (solution): 5/325 mg per 5 mL
oxycodone/ acetaminophen (Xartemis XR)	2 tablets every 12 hours	Swallow whole (do not crush, break, chew, cut, dissolve, or split) due to biphasic release system; swallow immediately Dose may be administered as early as 8 hours following the initial dose	Tablet: 7.5/325 mg
oxycodone/aspirin (Endodan, Percodan)	1 tablet every 6 hours	The maximum daily dose of aspirin should not exceed 4 grams or 12 tablets	Tablet: 4.8355 /325 mg
oxycodone/ ibuprofen	1 tablet per dose	Not to exceed a maximum of 4 tablets in 24 hours; do not exceed 7 days of therapy	Tablet: 5/400 mg
oxymorphone immediate-release (Opana)	10 to 20 mg every 4 to 6 hours	Administer on an empty stomach, at least 1 hour prior to or 2 hours after eating 5 mg dose is available for those with renal or hepatic impairment and for geriatric patients	Tablet: 5, 10 mg
pentazocine/ naloxone	1 to 2 tablets every 3 or 4 hours	Do not exceed 600 mg pentazocine per day	Tablet: 50/ 0.5 mg
tapentadol (Nucynta)	1 tablet every 4 hours	Doses greater than 700 mg on the first day and doses of greater than 600 mg on subsequent days are not recommended	Tablets: 50, 75,100 mg
tramadol (Ultram)	50 mg to 100 mg every 4 to 6 hours	Initiate at 25 mg every morning; titrate in 25 mg increments as separate doses every 3 days to reach 100 mg/day (25 mg four times daily), then the total daily dose may be increased by 50 mg as tolerated every 3 days to reach 200 mg/day (50 mg 4 times daily) After titration, tramadol 50 to 100 mg can be administered as needed for pain relief every 4 to 6 hours (not to exceed 400 mg per day)	Tablet: 50 mg



Drug	Starting Dose	Dosing Instructions	Available Strengths
tramadol/ acetaminophen (Ultracet)	to 6 hours	Not to exceed a maximum of 8 tablets in 24 hours; for the short-term (5 days or less) management of acute pain The elimination half-life of tramadol is increased in patients with severe renal impairment (CrCl < 30 mL/min), cirrhosis of the liver, or over 75 years, so the dosing interval should be extended	Tablets: 37.5/ 325 mg

Oxymorphone IR (Opana) should be given on an empty stomach; maximum concentration and area under the curve were increased 38% when given with a high-fat meal. Bioavailability of oxymorphone may also be increased in patients with hepatic or renal insufficiency. Formal studies have not yet been done. Label revisions to Lazanda dosage and administration provide an alternate titration strategy and modifications to the approved REMS. This is part of the Transmucosal Immediate-Release Fentanyl (TIRF) REMS Access Program.

Xartemis XR is an extended-release bilayer formulation of oxycodone and acetaminophen (immediateand extended-release layers) and is not interchangeable with other oxycodone/acetaminophen products due to differing pharmacokinetic profiles that affect the frequency of administration.

The only agent within this therapeutic class with abuse-deterrent properties is Oxaydo; Oxaydo contains sodium lauryl sulfate, inducing nasal passage irritation when crushed or snorted, and polyethylene oxide, forming a viscous mixture entrapping the opioid component to impede solvent extraction for intravenous abuse.^{253,254}



Opioid Morphine Equivalent Conversions²⁵⁵

This table is intended to provide an estimate of overall opioid exposure; it should not be used for dosing determinations (e.g., converting a patient from one opioid to another). Conversion factors may vary based on individual pharmacokinetics and duration of use (e.g., opioid-naïve versus chronic dosing). The same conversion is used for immediate- and extended-release oral products with the same opioid component unless otherwise specified. This table includes medications that are not reviewed in this class review for reference purposes. Likewise, some medications are not included in this table due to limited data.

Opioid	MME Conversion Factor
buprenorphine transdermal*	12.6
buprenorphine tablet or film	10
butorphanol	7
codeine	0.15
dihydrocodeine	0.25
fentanyl buccal, SL tablet, or lozenge [†]	0.13
fentanyl film or oral spray [†]	0.18
fentanyl nasal spray [†]	0.16
fentanyl patch [‡]	7.2
hydrocodone	1
hydromorphone	4
levorphanol tartrate	11
meperidine	0.1
methadone	3
morphine	1
nalbuphine	1
opium	1
oxycodone	1.5
oxymorphone	3
pentazocine	0.37
tapentadol	0.4
tramadol	0.1

^{*} Based on total micrograms exposure over 24 hours and assumes 1 mg parental buprenorphine = 75 mg oral morphine (e.g., 5 mcg/hr patch = 63 MME over 7 days = 9 MME/day).



[†] Multiply conversion factor by the number of micrograms in the dose.

[‡] Based on total micrograms exposure over 24 hours and assumes 1 mg parenteral fentanyl = 100 mg oral morphine (e.g., 25 mcg/hr patch = 180 MME over 3 days = 60 MME/day).

CLINICAL TRIALS

Search Strategies

Studies were identified through searches performed on PubMed and review of information sent by manufacturers. Search strategy included the FDA-approved use of all brand names in this class. Randomized, comparative, controlled trials performed in the United States comparing agents within this class in an outpatient setting for the approved indications are considered the most relevant in this category. Studies included for analysis in the review were published in English, performed with human participants, and randomly allocated participants to comparison groups. In addition, studies must contain clearly stated, predetermined outcome measure(s) of known or probable clinical importance, use data analysis techniques consistent with the study question, and include follow-up (endpoint assessment) of at least 80% of participants entering the investigation. Despite some inherent bias found in all studies, including those sponsored and/or funded by pharmaceutical manufacturers, the studies in this therapeutic class review were determined to have results or conclusions that do not suggest systematic error in their experimental study design. While the potential influence of manufacturer sponsorship and/or funding must be considered, the studies in this review have also been evaluated for validity and importance.

The following agents have demonstrated efficacy in placebo-controlled trials: fentanyl buccal tablet (Fentora), fentanyl nasal spray (Lazanda), fentanyl sublingual spray (Subsys), fentanyl sublingual tablet (Abstral), oxycodone/acetaminophen ER (Xartemis XR). ^{256,257,258,259,260,261}

butorphanol nasal spray and butalbital compound/codeine

In a double-blind, parallel-group study, patients with migraine (n=321) were randomly assigned to receive either butorphanol nasal spray 1 mg followed in 1 hour by an optional second 1 mg dose or butalbital compound with codeine administered orally (1 capsule containing butalbital 50 mg, caffeine 40 mg, aspirin 325 mg, and codeine phosphate 30 mg). Patients were instructed to self-administer medication when migraine pain reached intensity of moderate or severe and to record study-related events in a diary for 24 hours post-treatment. Efficacy analyses were performed on data from 275 patients who received study medication and returned a patient diary. During the first 2 hours after treatment, butorphanol was more effective than butalbital compound/codeine in treating migraine pain as measured by pain intensity difference scores, percentage of responders (pain decreased to mild or none), percentage of pain-free patients, and degree of pain relief, with a more rapid time to onset of 15 minutes. A similar percentage of patients in the 2 groups used rescue medication during the first 4 hours, after which more butorphanol-treated than butalbital compound/codeine-treated patients used rescue medication. Butorphanol patients had more side effects, less improvement in digestive symptoms, and less improvement in functional ability than butalbital compound/codeine patients.

fentanyl oral transmucosal (Actiq) and morphine IR

In a randomized, double-blind, cross-over trial with 134 adult ambulatory cancer patients, fentanyl oral transmucosal and morphine sulfate immediate-release (MSIR) were compared for the management of breakthrough pain. ²⁶³ Enrolled patients were stabilized on a fixed schedule opioid regimen of either morphine sulfate or transdermal fentanyl and an effective MSIR dose of 15 to 60 mg up to 4 times daily for breakthrough pain. In an open-label fashion, fentanyl oral transmucosal was administered to establish the effective dose for breakthrough pain for 69% of patients. Double-blind randomization



occurred and then a set of capsules and oral transmucosal delivery systems (1 placebo unit per set being either capsule or transmucosal unit) were administered for each breakthrough pain dosing. During the blinded study, fentanyl oral transmucosal was significantly better than MSIR for pain intensity reduction, pain relief, and pain intensity differences. Patients favored the fentanyl oral transmucosal for breakthrough pain based on global performance.

oxymorphone IR (Opana) and oxycodone IR

In a double-blind, parallel-group study, oxymorphone IR was compared with placebo for efficacy and with oxycodone IR and placebo for safety in patients with acute moderate to severe post surgical pain. Three hundred patients received oxymorphone IR 10, 20, or 30 mg; oxycodone IR 10 mg; or placebo. All oxymorphone IR doses were superior to placebo for providing pain relief for 8 hours (p<0.05), each with a significant analgesic dose response compared to placebo (p<0.001). All oxymorphone IR groups maintained analgesia for 48 hours. The median dosing interval was over 9.5 hours for oxymorphone IR 30 mg. Opioid-related adverse events, similar among groups, were generally mild or moderate; the overall safety profile was comparable to that of oxycodone IR.

oxycodone/ibuprofen (Combunox) versus oxycodone/acetaminophen (Percocet) versus hydrocodone/acetaminophen (Vicodin ES)

In a multicenter, randomized, double-blind, placebo- and active-controlled, parallel-group, single-dose study, patients experiencing moderate to severe pain after surgical removal of 2 or more ipsilateral impacted third molars were randomly assigned to receive oxycodone/ibuprofen 5/400 mg, oxycodone/acetaminophen 5/325 mg, hydrocodone/acetaminophen 7.5/500 mg, or placebo. The primary outcome measures were total pain relief through 6 hours after dosing, sum of pain intensity differences through 6 hours (SPID6), and adverse events. Oxycodone/ibuprofen 5/400 mg provided significantly greater analgesia 6 hours after dosing compared with oxycodone/acetaminophen 5/325 mg, hydrocodone/acetaminophen 7.5/500 mg, and placebo (p<0.001, oxycodone/ibuprofen 5/400 mg versus all other treatments). Values for SPID6 also differed significantly for oxycodone/ibuprofen 5/400 mg compared with all other treatments (p<0.001). Oxycodone/ibuprofen 5/400 mg was significantly more effective compared with the other treatments on all secondary endpoints (p<0.001), with the exception of the time to onset of analgesia. The lowest frequency of nausea and vomiting occurred in the groups that received oxycodone/ibuprofen 5/400 mg (6.5% and 3.2%, respectively) and placebo (3.2% and 1.6%).

oxycodone/ibuprofen (Combunox) versus oxycodone (Roxicodone) versus ibuprofen (Motrin)

In a multicenter, randomized, double-blind, placebo- and active-controlled, parallel-group trial, women experiencing moderate to severe pain between 14 and 48 hours after surgery were randomized to receive a single dose of oxycodone/ibuprofen, ibuprofen, oxycodone, and placebo. ²⁶⁶ Four hundred fifty-six women participated in the study. Combination treatment was associated with significantly better scores for total pain relief 6 hours after dosing and sum of pain intensity differences 6 hours after dosing compared with ibuprofen alone (p<0.02 and p<0.015, respectively), oxycodone alone (p<0.009 and p<0.001), or placebo (both p<0.001). Fewer patients receiving combination treatment required rescue medication, and the time to use of rescue medication was significantly longer in the combination treatment group compared with the other groups (p<0.05). The onset of pain relief



occurred within 15 minutes of dosing with all regimens. Nausea was the most frequently reported adverse event in all groups, highest with placebo and followed by oxycodone, ibuprofen, and combination treatment.

In the multicenter, double-blind, double-dummy, parallel-group investigation, 498 patients with moderate to severe pain within 5 hours after extraction of 2 or more impacted third molars were randomized to single doses of oxycodone/ibuprofen 5/400 mg, ibuprofen 400 mg, oxycodone 5 mg, or placebo. Combination therapy was associated with greater analgesia than ibuprofen alone, oxycodone alone, or placebo, as measured by the sum of pain intensity difference over 6 hours (p<0.001 versus oxycodone or placebo, p=0.002 versus ibuprofen) and total pain relief through 6 hours (p<0.001 versus oxycodone or placebo, p=0.012 versus ibuprofen). Combination therapy was well tolerated, and pharmacokinetic evaluation implied no interaction between oxycodone and ibuprofen.

oxymorphone (Opana) versus oxycodone IR versus placebo

A multicenter, randomized, double-blind, active- and placebo-controlled, parallel-group study was conducted in men and women aged 18 years and older undergoing abdominal surgery. Patients were randomized to receive oxymorphone 10 or 20 mg, oxycodone 15 mg, or placebo every 4 to 6 hours. The study included single-dose and 48-hour efficacy assessments. The primary efficacy endpoint was the median time to study discontinuation for all causes. Three hundred thirty-one patients were included in the study. The median time to study discontinuation was significantly longer for all active treatments compared with placebo (oxymorphone 10 mg, 17.9 hours; oxymorphone 20 mg, 20.3 hours; oxycodone 15 mg, 24.1 hours; placebo, 4.8 hours; p<0.006). Oxymorphone 20 mg was significantly more effective than placebo over the 6-hour single-dose evaluation (p<0.05). With multiple dosing, all active-treatment groups had significantly lower least squares mean current and average pain intensities compared with placebo (p<0.004 and p<0.005, respectively). Discontinuations due to treatment-emergent adverse events did not differ significantly among the groups.

tapentadol (Nucynta) versus morphine IR

Patients (n=400) undergoing molar extraction were randomized to receive single doses of tapentadol 25, 50, 75, 100, or 200 mg, morphine sulfate 60 mg, ibuprofen 400 mg, or placebo. Hean total pain relief over 8 hours (TOTPAR-8) was the primary endpoint. Secondary endpoints included mean total pain relief over 4 hours (TOTPAR-4) and onset of analgesia. Of all measured endpoints, only mean TOTPAR-4 was higher (and onset of action appeared more rapid) for tapentadol 200 mg than morphine sulfate 60 mg. Pain relief scores with morphine sulfate 60 mg were between those of tapentadol 100 and 200 mg. The incidence of nausea and vomiting appeared to be lower with all doses of tapentadol compared with morphine sulfate 60 mg but was not statistically significant.

tapentadol (Nucynta) versus oxycodone (Roxicodone)

A 10-day, phase 3, randomized, double-blind, active- and placebo-controlled study compared the efficacy and tolerability of tapentadol, oxycodone, and placebo in 666 patients with uncontrolled osteoarthritis pain who were candidates for primary replacement of the hip or knee as a result of end-stage degenerative joint disease. Patients received tapentadol 50 mg or 75 mg, oxycodone 10 mg, or placebo every 4 to 6 hours while awake. The primary endpoint was the SPID over 5 days. Prespecified noninferiority comparisons with oxycodone were performed with respect to efficacy and tolerability. Five-day SPID was significantly lower in those treated with tapentadol or oxycodone (all p<0.001).



Tapentadol 50 and 75 mg and oxycodone 10 mg were associated with significant reductions in pain intensity compared with placebo based on 2- and 10-day SPID, as well (all p<0.001). The efficacy of tapentadol 50 and 75 mg was noninferior to that of oxycodone 10 mg; however, the incidence of nausea, vomiting, and constipation was significantly lower for both doses of tapentadol compared with oxycodone (p<0.001).

tramadol/acetaminophen (Ultracet) versus tramadol (Ultram)

A total of 456 patients with moderate to severe pain within 5 hours of extraction of 2 or more third molars were randomized to receive 2 identical encapsulated tablets containing tramadol/acetaminophen 37.5/325 mg, tramadol 50 mg, or placebo. Tramadol/acetaminophen was superior to tramadol (p<0.001) or placebo (p<0.001) on all efficacy measures, including total pain relief over 6 hours, sum of pain intensity differences, and sum of both. The most common adverse events with active treatment were nausea, dizziness, and vomiting, which occurred more frequently in the tramadol group than in the tramadol/acetaminophen group.

tramadol/acetaminophen (Ultracet) versus codeine/acetaminophen

A randomized, double-blind, parallel-group, active-control, double-dummy trial compared the efficacy and tolerability of tramadol/acetaminophen 37.5 /325 mg tablets with codeine/acetaminophen capsules 30 /300 mg in 462 patients with chronic nonmalignant low back pain, osteoarthritis, or both. Pain intensity was assessed hourly for 6 hours each week over a 4-week period. Pain relief and changes in pain intensity were comparable in both groups throughout the study. Equivalent mean doses and maximum daily doses used in each group were similar. The overall incidence of adverse events was comparable, with more patients in the codeine/acetaminophen group reporting somnolence (24%versus 17%, p=0.05) and constipation (21% versus 11%, p<0.01) than the tramadol/acetaminophen group.

A multicenter, randomized, double-blind, active- and placebo-controlled trial evaluated tramadol plus acetaminophen for orthopedic and abdominal post surgical pain.²⁷³ Patients with moderate pain or greater were randomized to an initial 2 tablets of 37.5 mg tramadol plus 325 mg acetaminophen (n=98), codeine 30 mg plus acetaminophen 300 mg (n=109), or placebo (n=98). Thereafter, they received 1 to 2 tablets every 4 to 6 hours, as needed for pain, for 6 days. Tramadol plus acetaminophen was superior to placebo for total pain relief, sum of pain intensity differences, and sum of pain relief and pain intensity differences (p≤0.015). For average daily pain relief, average daily pain intensity, and overall medication assessment, tramadol plus acetaminophen was superior to placebo (p≤0.038); codeine plus acetaminophen did not separate from tramadol plus acetaminophen in any criteria. Discontinuation because of adverse events occurred in 8.2% of tramadol plus acetaminophen, 10.1% of codeine plus acetaminophen, and 3% of placebo patients. Except for constipation and vomiting being more prevalent in codeine plus acetaminophen patients, adverse events were similar for active treatments.

A 4-week, randomized, double-blind, parallel-group, multicenter trial compared tramadol/acetaminophen 37.5 /325 mg with codeine/acetaminophen 30/300 mg for the management of chronic nonmalignant low back pain, osteoarthritis pain, or both in 462 adults. Pain relief (scale, 0 = none to 4 = complete) and pain intensity (scale, 0 = none to 3 = severe) were measured after 30 minutes and then hourly for 6 hours after the first daily dose each week. Pain relief and changes in pain intensity were comparable from Day 1 and lasted for at least 6 hours. Total pain relief scores and sum



of pain intensity differences were also comparable throughout. Overall assessments of safety and efficacy by patients and investigators were similar for the 2 treatment groups.

tramadol/acetaminophen (Ultracet) versus hydrocodone/acetaminophen (Vicodin)

In a single-center, double-blind, parallel-group, placebo- and active-controlled study in adults with at least moderate pain after extraction of 2 or more impacted third molars, patients were randomized to receive 1 to 2 tramadol/acetaminophen 37.5 /325 mg tablets, 1 hydrocodone/acetaminophen 10 /650 mg tablet, or placebo.²⁷⁵ Two hundred adults took part in the study. The median time to onset of pain relief was approximately 34 minutes with tramadol/acetaminophen tablets and 25.4 minutes with hydrocodone/acetaminophen. Although the median time to onset of pain relief was shorter with hydrocodone/acetaminophen, 2 tramadol/acetaminophen tablets had comparable efficacy to hydrocodone/acetaminophen. The median time to re-medication with a supplemental analgesic agent was 169 minutes in the tramadol/acetaminophen group and 204 minutes in the hydrocodone/acetaminophen group; however, the duration of pain relief was not significantly different between the groups. The overall incidence of adverse events was lower with tramadol/acetaminophen (0%) than with hydrocodone/acetaminophen (4%) or placebo (10%).

META-ANALYSES

Meta-analyses evaluating single-agents within this class have been published, but meta-analyses comparing agents within this class are limited. 276,277,278,279,280,281,282 In addition, some meta-analyses do not differentiate short-acting and long-acting opioids.

A few meta-analyses have compared agents within this class in patients with breakthrough or general cancer pain. One meta-analysis of 10 randomized clinical trials for breakthrough cancer pain compared various forms of fentanyl (e.g., nasal spray, sublingual tablets, buccal film, transmucosal) to morphine sulfate immediate-release in pain intensity difference compared to placebo up to 60 minutes following intake. Most fentanyl formulations, excluding sublingual tablets, resulting in a greater pain intensity difference compared to placebo at 15 minutes following intake while all fentanyl formulations showed a difference at 30 minutes. However, morphine sulfate did not demonstrate a difference in pain intensity until 45 minutes. Likewise, only nasal fentanyl spray produced a clinically meaningful difference (pain intensity change ≥ 2) at 15 minutes. An earlier meta-analysis of 5 trials found similar results. Page 284

A meta-analysis of 14 studies (n=3,521) assessed various analgesic combinations (e.g., acetaminophen with codeine [various strength combinations], acetaminophen with hydrocodone [various strength combinations], non-opioids, codeine/butalbital/aspirin/caffeine, oxycodone with ibuprofen, and ibuprofen with codeine) for postoperative pain following third-molar surgery. Of all combinations, ibuprofen 400 mg in combination with oxycodone 5 mg had superior efficacy in sum of pain intensity at 6 hours scores (6.44; range of all agents, 1.46 to 6.44) and total pain relief at 6 hours scores (9.31; range of all agents, 3.24 to 10.3).

A Cochrane review of 35 other Cochrane reviews (approximately 45,000 participants in approximately 350 studies) evaluated single-dose analgesics for acute postoperative pain in adults, including non-opioid analgesics and dental surgeries. The primary outcome assessed was at least 50% pain relief over 4 to 6 hours compared to placebo. The authors calculated number-needed-to-treat (NNT) in reliable studies to achieve this primary outcome calculated for all agents and ranged from 1.5 to 20



among all agents. NNTs of agents in this class were 2.2 (95% CI, 2.3 to 3.3), 3.9 (95% CI, 3.3 to 4.7), 2.7 (95% CI, 2.4 to 3.1), and 1.8 (95% CI, 1.6 to 2.2), for codeine 60 mg with acetaminophen 800 to 1,000 mg, codeine 60 mg, codeine 60 mg with acetaminophen 600 to 650 mg, oxycodone/acetaminophen 10/650 mg, and oxycodone/acetaminophen 10/1,000 mg, respectively.

SUMMARY

Pain management must be individualized for each patient. There are many equally effective opioid analgesic products available, differing in specific opioid (and co-analgesics), dosage form, and duration of action. Many are available in clinically effective generic forms, including combinations of non-narcotic acetaminophen, aspirin, or ibuprofen with the opioids hydrocodone or oxycodone. Although some manufacturers market unique strengths of these combination agents, the minor changes in the doses of acetaminophen, ibuprofen, and/or opioid in these products have not been shown to offer any advantage over similar generic combinations. Similarly, there are no data to suggest that a particular formulation of fentanyl (Abstral, Actiq, Fentora, Lazanda, Subsys) is safer or more effective for breakthrough cancer pain.

Dihydrocodeine/caffeine/acetaminophen (Trezix), tapentadol (Nucynta), tramadol/acetaminophen (Ultracet), and oxymorphone (Opana) have not shown increased efficacy when compared to other opioids.

Oxaydo is an immediate-release opioid analgesic with abuse-deterrent properties intended to discourage abuse of the medication. These preventative measures offer no analgesic advantage over existing products. While it is acknowledged that diversion and misuse of opioids may be commonplace, patients should be evaluated to determine whether such preventative measures are required.

Xartemis XR is the first extended-release oral combination of oxycodone and acetaminophen approved for acute pain severe enough to require opioid treatment and for which alternative treatment options are inadequate.

Clinical guidelines do not recommend one opioid agent over another.

All agents within this class are considered Controlled Substances and contain a box warning regarding serious risks of misuse, abuse, addiction, overdose, and death; fentanyl, hydrocodone, hydromorphone, levorphanol, meperidine, morphine, oxycodone, oxymorphone, and tapentadol containing products and codeine tablets are Schedule II; codeine and dihydrocodeine combination products are Schedule III; and butorphanol, pentazocine, and tramadol containing products are Schedule IV.

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